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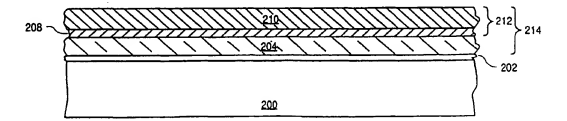
#### **EUROPEAN PATENT APPLICATION**

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- (54) Method of forming tungsten-silicide
- (57) A method of forming a silicide on a silicon layer. First, a monosilane based tungsten-silicide layer is

formed on the silicon layer. Next, a dichlorosilane based tungsten-silicide layer is formed on the monosilane based tungsten-silicide layer.



#### FIG. 2D

#### Description

The present invention relates to the field of thin films and their methods of manufacturing and more specifically, to composite silicon/tungsten-silicide thin films and their methods of fabrication.

As features of VLSI circuits continue to shrink, the necessity of decreasing the resistance associated with interconnection paths and gate electrodes becomes ever more pressing. Polysilicon has been widely used for many years in the formation of gate electrodes and interconnects. Polysilicon, however, has been somewhat limited by its high bulk resistivity of 0.7 mohm-cm. Polycide films, which consist of a low resistance silicide formed on top of a doped polysilicon film, have been used in place of polysilicon films in an attempt to further decrease the resistance of gate electrodes and interconnects to thereby create faster and lower power consuming integrated circuits.

A polycide film comprising a tungsten-silicide layer formed on a doped polysilicon layer has been widely used as gate electrodes and interconnections in many metal oxide semiconductor (MOS) processes. In the fabrication of a doped polysilicon/tungsten-silicide gate electrode, as shown in Figure 1a, a doped polysilicon layer 104 of approximately 1000 angstroms (Å) is blanket deposited onto a gate dielectric layer 102 formed over the surface of a wafer or substrate 100. A tungstensilicide layer 106, of the same order of magnitude in thickness as polysilicon layer 104 is then deposited over doped polysilicon layer 104 to form a "polycide" stack 108. Tungsten-silicide layer 106 can be formed by chemical vapor deposition (CVD) utilizing a reduction reaction of monosilane (SiH<sub>4</sub>) and tungsten hexafluorine (WFs). Such a deposition technique forms a good quality film having a uniform thickness across the surface of the wafer and a relatively smooth top surface. Unfortunately, however, utilizing a SiH4 reduction of WF<sub>6</sub> to form tungsten-silicide layer 106 incorporates a substantial amount of fluorine atoms into the tungstensilicide layer which can later damage or destroy fabricated devices.

In order to reduce fluorine incorporation, tungstensilicide layer 110 can be formed by CVD utilizing a reduction reaction of dichlorosilane (DCS or SiH2Cl2) and tungsten hexafluorine (WF6). Although such a DCS based reaction substantially eliminates fluorine incorporation, the film is of low quality because dopant impurities in polysilicon layer 104 prevent good nucleation from occurring. As a result, as shown in Figure 1b, the film forms nonuniformly over the surface of the wafer and has an unacceptably rough surface. Such a polycide film 112 is unacceptable for VLSI manufacturing. Another problem associated with forming a DCS based tungsten-silicide layer on a doped polysilicon layer is that such a deposition technique can form a tungsten rich interface which is known to negatively impact device performance and cause reliability problems such as

electromigration and high film stress. The presence of polysilicon dopants at the interface can also lead to the formation of an undesired tungsten rich interface.

It has been discovered that in order to produce a tungsten-silicide film of good quality by a DCS reduction of WF6, it is necessary to form a thin, at least 150Å, undoped polysilicon capping layer 114 on doped polysilicon layer 104. Undoped polysilicon capping layer 114 provides a barrier between the dopant impurities in doped polysilicon layer 104 and DCS based tungstensilicide film 116. In this way, a DCS based tungsten-silicide film 116 can be formed with good uniformity and a substantially smooth surface across a wafer. Unfortunately, however, such a technique is only viable for polycide films 120 having a thickness greater than 1000Å. For thinner films, less than 1000Å, the undoped capping layer 114, which must still be at least 150Å to produce a sufficient barrier, comprises too much of the total film thickness and results in a substantial reduction in the total dopant concentration of the polycide film 120 which in turn results in an increase in total resistance. Attempts to further increase the doping concentration level of the polysilicon layer 104 to offset capping layer 114 results in dopant diffusion into gate oxide 102 and consequently to device performance degradation and damage.

Thus, what is desired is a method of forming a thin low resistance polycide film having a uniform thickness and a smooth top surface which does not incorporate an undesirable amount of fluorine, and which does not have a tungsten rich interface.

Methods of forming a silicide on a doped silicon layer, without an undoped silicon cap are described. In a first method, a doped silicon layer is exposed to a first gas mix comprising monosilane (SiH<sub>4</sub>) to form a monosilane exposed doped silicon surface. Next, a first tungsten-silicide layer, formed by a reduction reaction of SiH<sub>4</sub> and WF<sub>6</sub>, is formed on the monosilane exposed silicon surface. A second tungsten-silicide layer, formed by a reduction reaction of SiH<sub>2</sub>Cl<sub>2</sub> and WF<sub>6</sub>, is then formed on the first tungsten-silicide layer. In a second method, a doped amorphous silicon layer is formed. A tungsten-silicide layer, formed by a reduction reaction of SiH<sub>2</sub>Cl<sub>2</sub> and WF<sub>6</sub>, is then deposited onto the amorphous silicon layer.

Figure 1a is an illustration of a cross-sectional view showing a polycide film comprising a doped polysilicon layer and a tungsten-silicide layer formed from SiH<sub>4</sub> and WF<sub>6</sub>.

Figure 1b is an illustration of a cross-sectional view showing a polycide film comprising a doped polysilicon layer and a tungsten-silicide layer formed from SiH<sub>2</sub>Cl<sub>2</sub>.

Figure 1c is an illustration of a cross-sectional view showing a polycide film comprising a doped polysilicon layer, an undoped polysilicon capping layer, and a tungsten-silicide layer formed from SiH<sub>2</sub>Cl<sub>2</sub> and WF<sub>6</sub>.

Figure 2a is an illustration of a cross-sectional view showing a gate dielectric layer formed on a substrate and a doped polysilicon layer formed on the gate die-

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Figure 2b is an illustration of a cross-sectional view showing a monosilane pretreatment of the polysilicon layer on the substrate of Figure 2a.

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Figure 2c is an illustration of a cross-sectional view showing the formation of a tungsten-silicide layer from a reaction of SiH<sub>4</sub> and WF<sub>6</sub> on the substrate of Figure 2h

Figure 2d is an illustration of a cross-sectional view showing the formation of a tungsten-silicide layer from a reaction of  $SiH_2Cl_2$  and  $WF_6$  on the substrate of Figure 2c.

Figure 3a is an illustration of a cross-sectional view showing the formation of an amorphous silicon layer on a substrate.

Figure 3b is an illustration of a cross-sectional view showing the formation of a dichlorosilane based tungsten-silicide layer on the substrate of Figure 3a.

Novel silicon/tungsten-silicide composite thin films and their methods of fabrication are described. In the following description, numerous specific details are set forth, such as specific materials, thicknesses, and process parameters, in order to provide a thorough understanding of the present invention. It will be obvious, however, to one skilled in the art that the present invention may be practiced without these specific details. In other instances, other well known semiconductor processes and machinery have not been described in particular detail in order to avoid unnecessarily obscuring the present invention.

The first embodiment of the present invention describes a novel method of forming a novel composite film comprising a lower silicon layer, an interfacial tungsten-silicide layer formed by chemical vapor deposition (CVD) utilizing a process gas mix of monosilane (SiH<sub>4</sub>) and tungsten hexafluorine (WFs), and a top tungstensilicide layer formed by CVD utilizing a process gas mix comprising dichlorosilane (DCS or SiH2Cl2) and WF6. The novel composite film of the present invention can be formed thin (less than 1000Å), and uniformly over the entire surface of a wafer. The film is characterized by a smooth top surface and a low sheet resistance. The composite film of the present invention is ideally suited for use as a gate electrode or as an interconnection in an integrated circuit such as a microprocessor or memory device, but may be used in other polycide applications, if desired.

The present invention will be explained with respect to the formation of a polycide film to be used as a gate electrode in a metal oxide semiconductor (MOS) integrated circuit. In such a process, a semiconductor substrate or wafer 200 is provided. It is to be appreciated that substrate 200 is preferably a single crystalline silicon substrate doped to a suitable concentration level and may include well known wells and field isolation regions and other conventional features. A gate dielectric layer 202, such as SiO<sub>2</sub>, having a thickness less than 100Å, is formed over substrate 200.

Next, as shown in Figure 2a, a silicon layer 204 is formed over gate dielectric layer 202. According to the preferred embodiment of the present invention, silicon layer 204 is a polycrystalline silicon layer (polysilicon) doped with impurities, such as phosphorous, to a concentration level between 1x10<sup>20</sup>/cm<sup>3</sup>-6x10<sup>20</sup>/cm<sup>3</sup>. Silicon layer 204 is preferably a polycrystalline silicon layer formed to a thickness of approximately 500Å or less (for a polycide film of 1000Å or less) by chemical vapor deposition (CVD) in a separate chamber of the same cluster tool used to deposit the subsequently deposited tungsten-silicide layers described below.

It is to be appreciated that according to the first embodiment of the present invention, silicon layer 204 need not necessarily be a polysilicon layer, but may be any other type of silicon layer, such as amorphous silicon, if desired. Additionally, silicon layer 204 need not necessarily be a highly doped silicon layer and may be a low doped or undoped layer, if desired. It is to be appreciated, however, that an advantage of the present invention is its ability to form a uniform, high quality, tungsten-silicide film on a doped silicon layer. However, any well known silicon layer and method of fabrication may be used to form silicon layer 204.

Next, according to the first preferred embodiment of the present invention, substrate or wafer 200 is transferred from the polysilicon deposition chamber into a CVD tungsten-silicide deposition chamber such as available in a Centura high temperature film (HTF) tool available from Applied Materials of Santa Clara, California. The tungsten-silicide deposition chamber is preferably lamp heated and preferably has a gas delivery system through the lid with a susceptor to shower head spacing of approximately 475 mills. Additionally, the tungsten-silicide deposition chamber is preferably a single wafer chamber capable of being evacuated to less than 0.2 T. Additionally, the gas delivery system is coupled to and can supply sufficient volumes of the following process gases: SiH<sub>4</sub>, SiH<sub>2</sub>Cl<sub>2</sub>, and WF<sub>6</sub>, and is coupled to and can supply sufficient volumes of an inert gas or gases such as argon (Ar), helium (He), and nitrogen  $(N_2)$ .

Next, as shown in Figure 2b, polysilicon layer 204 is preconditioned by exposing polysilicon layer 204 to a gas comprising monosilane (SiH<sub>4</sub>). The goal of the monosilane pretreatment (or "monosilane soak") is to form a uniform layer (tens of Angstroms) of monosilane 206 across the surface of polysilicon layer 204 prior to flowing a source of tungsten into the deposition chamber. In this way, when WF<sub>6</sub> is subsequently flowed into the deposition chamber, a uniform layer of monosilane is already provided across the surface of the polysilicon layer 204 allowing for a uniform deposition of tungsten-silicide on polysilicon layer 204. Although preconditioning polysilicon layer 204 is desired to help achieve good tungsten-silicide deposition, it is not required to practice the present invention.

Preconditioning of polysilicon layer 204 can be ac-

complished by flowing between 100-1000 sccm of monosilane, with 300 sccm being preferred, into the deposition chamber, at a temperature between 450-625°C with 550°C being preferred, for a time between 5-60 seconds with 40 seconds being preferred. The preconditioning step occurs at a pressure between 0.5-15 T with one T being preferred. It is to be appreciated that preconditioning polysilicon layer 204 at too high of a pressure can cause undesirable silicon deposition and subsequent tungsten-silicide deposition on chamber walls and windows and on the backside of the susceptor. Argon can be included in the monosilane preconditioning gas mix in order to help prevent undesired silicon formation in the chamber.

Next, as shown in Figure 2c, a thin monosilane (SiH<sub>4</sub>) based tungsten-silicide layer 208 is formed onto polysilicon layer 204. The purpose of monosilane based tungsten-silicide layer 208 is to provide an interfacial layer which has a uniform thickness, a smooth surface, and a consistent resistivity across the surface of wafer 200. Tungsten-silicide layer 208 is formed by chemical vapor deposition (CVD) utilizing a reduction reaction of monosilane (SiH<sub>4</sub>) and WF<sub>6</sub>. Monosilane based tungsten-silicide layer 208 is deposited to a thickness between 10-500Å with 100Å being preferred. Tungsten-silicide layer 208 is formed thick enough to uniformly form at least one or two monolayers of tungsten-silicide onto polysilicon layer 204 but yet is kept thin enough so as to limit the amount of fluorine incorporated into the deposited film. Because tungsten-silicide layer 208 is formed onto a monosilane exposed polysilicon surface 206, a thin uniform layer of tungsten-silicide 208 can be reliably formed across the surface of wafer 200.

Monosilane based tungsten-silicide layer 208 is preferably deposited insitu (i.e. in the same chamber and immediately after) the monosilane preconditioning step. Tungsten-silicide layer 208 is preferably formed by introducing a gas mix comprising approximately four sccm of WF<sub>6</sub> and approximately 300 sccm of monosilane into a reaction chamber for approximately two-three seconds at a pressure between 0.2-10 T with one-two T being preferred and at a temperature between 450-625°C with 550°C being preferred. An inert gas or gases such as argon, helium and nitrogen can be included into the gas mix in order to prevent undesired tungsten-silicide deposition onto the reaction chamber and to provide a carrier for WF<sub>6</sub>, if desired.

It is to be appreciated that the reaction of SiH<sub>4</sub> and WF<sub>6</sub> produces tungsten-silicide (WSi<sub>x</sub>) where X≥2.0. As such, the resistivity of the deposited tungsten-silicide film 208 can be tailored to specific needs by adjusting the gas ratio of SiH<sub>4</sub>:WF<sub>6</sub>. The greater the flow of SiH<sub>4</sub> relative to WF<sub>6</sub>, the greater will be the silicon content in the tungsten-silicide film. The greater the silicon content in the film, the higher will be the resistivity of the film. Additionally, tungsten-silicide layer 208 is preferably deposited so that it has a flat Si:W composition ratio (i.e. the Si:W ratio is substantially the same throughout the

thickness of the film).

Next, as shown in Figure 2d, a dichlorosilane (DCS or SiH<sub>2</sub>Cl<sub>2</sub>) based tungsten-silicide layer 210 is formed onto monosilane based tungsten-silicide layer 208. Tungsten-silicide layer 210 is formed by chemical vapor deposition utilizing a reduction reaction of WF<sub>6</sub> and DCS and is preferably formed insitu (i.e. in the same chamber and immediately after) monosilane based tungsten-silicide layer 208. In order to reduce the fluorine content in the polycide film, DCS based tungsten-silicide layer 210 makes up the bulk of the dual layer tungsten-silicide film 212. DCS based tungsten-silicide layer 210 is preferably formed to a thickness between 300-700Å for a 1000Å polycide film. For a polycide stack greater than 1000Å, DCS based tungsten-silicide layer 210 should preferably be formed to a thickness greater than 500Å.

The gas flow ratio of DCS:WF<sub>6</sub> can be chosen to obtain a desired tungsten-silicide composition ratio and therefore a desired resistivity of tungsten-silicide film 210. Because DCS based tungsten-silicide film 210 comprises the bulk of the dual layer tungsten film 212, it must be formed to a thickness and with the appropriate Si:W composition ratio to obtain the desired resistivity and thickness for polycide film 214.

DCS based tungsten-silicide layer 210 can be formed by flowing a gas mix comprising approximately 200 sccm of DCS and approximately 4 sccm of WF<sub>6</sub> into a reaction chamber at a pressure between 0.2-10 T with approximately one T being preferred, and at a temperature between 450-625°C with 550°C being preferred. An inert gas or gases, such as argon, helium and nitrogen, can be included in the gas mix in order to prevent undesired tungsten-silicide deposition onto the reaction chamber and to provide a carrier for WF<sub>6</sub>, if desired. It is to be appreciated that DCS based tungsten-silicide film 210 is able to be formed uniformly and with a smooth surface because interfacial monosilane based tungsten-silicide layer 208 prevents interaction with doped polysilicon layer 204.

It is to be noted that it is undesirable to have a tungsten rich interface between dual layer tungsten-silicide film 212 and polysiticon layer 204. A tungsten rich interface can cause electromigration which can lead to void formation which in turn can cause open circuits. Additionally, a tungsten-rich interface can result in a high film stress which can lead to peeling. As such, according to the first preferred embodiment of the present invention, deposition parameters are chosen so as to form a monosilane based tungsten-silicide layer 208 with a Si:W content ratio greater than or equal to the Si:W content ratio of the DCS based tungsten-silicide layer 210. That is, monosilane based tungsten-silicide layer 208 is preferably the same as or more silicon rich than DCS based tungsten-silicide layer 210.

After deposition of DCS based tungsten-silicide layer 210, a "monosilane cap" can be formed on DCS based tungsten-silicide layer 210, if desired. Flowing monosilane over tungsten-silicide layer 210 allows any

loose tungsten bonds to react with monosilane and form tungsten-silicide. Forming a "monosilane cap" helps reduce stress in the DCS based tungsten-silicide layer. A "monosilane cap" can be formed insitu by flowing approximately 200 scan of monosilane for approximately 15 seconds into the tungsten-silicide deposition chamber at a pressure of approximately one T and at a temperature of approximately 550°C.

After formation of a "monosilane cap," substrate 200 is removed from the tungsten-silicide deposition chamber. Substrate 200 is then eventually annealed to activate and drive dopants in polysilicon layer 204 and to alter the stoichiometry of tungsten-silicide films 208 and 210, from WSi<sub>x</sub> X≥2.0≤3.5 to WSi<sub>x</sub> X≈2.0 which reduces the sheet resistance of the polycide film. Any well known anneal method and apparatus can be used to anneal substrate 200, such as a Rapid Thermal Anneal (RTA), at approximately 1000°C, in a nitrogen ambient, for a time between 30-60 seconds, or a standard furnace anneal at 850°C for 30 minutes.

At this point, the formation of a novel polycide film comprising a doped polysilicon layer 204 and interfacial monosilane based tungsten-silicide layer 208, and a bulk DCS based tungsten-silicide layer 210 is now complete. The low resistance monosilane based tungsten-silicide interfacial layer of the first embodiment of the present invention enables a thin (less than 1000Å), low resistance (600-1500  $\mu$   $\Omega$ -cm) polycide film to be formed uniformly across the surface of a wafer without substantial fluorine incorporation therein. Conventional photolithographic and etching techniques can be used to pattern polycide film 214 into a plurality of gate electrodes. Well known doping and interconnection techniques can then be utilized to complete the fabrication of an integrated circuit.

A second embodiment of the present invention describes a novel composite film comprising a lower amorphous silicon layer and a top tungsten-silicide layer formed by CVD utilizing a process gas mix comprising dichlorosilane (DCS) and tungsten hexafluorine (WF<sub>6</sub>). The novel composite film of the second preferred embodiment of the present invention can be formed thin (less than 1000Å) and uniformly across the surface of a wafer. The thin film is characterized as having a smooth top surface and a low sheet resistance. Additionally, the film can be formed without undesirable fluorine incorporation.

According to the second method of the present invention, an amorphous silicon layer 304 is blanket deposited onto gate dielectric layer 202 formed on substrate 200. Amorphous layer 304 is preferably formed to a thickness of less than 500Å by chemical vapor deposition (CVD) and is insitu doped with impurities, such as phosphorous, to a concentration level between 1x10<sup>20</sup>/cm³ to 6x10<sup>20</sup>/cm³. Amorphous silicon layer 304 can be formed by CVD by flowing monosilane (SiH<sub>4</sub>), H<sub>2</sub>, and a dopant source, such as phosphine, into a reaction chamber at a pressure between 3-200T and at a

temperature between 600-630°C. Any well known method, however, can be utilized to form amorphous silicon layer 304.

Next, as shown in Figure 3b, substrate 200 is transferred into a tungsten-silicide deposition chamber such as described above. A dichlorosilane (SiH $_2$ Cl $_2$  or DCS) based tungsten-silicide layer 306 is then deposited onto amorphous silicon layer 304. DCS based tungsten-silicide layer is formed by chemical vapor deposition utilizing a reduction reaction of WF $_6$  and DCS. The gas flow ratio of DCS:WF $_6$  is chosen to obtain a desired Si:W composition ratio and therefore the desired resistivity of tungsten-silicide film 306. DCS based tungsten-silicide 306 is preferably formed so as to generate has a flat Si:W composition ratio.

DCS based tungsten-silicide layer 306 is preferably deposited to a thickness less than 500Å by flowing a gas mix comprising approximately 200 scan of DCS and approximately four sccm of WF<sub>6</sub> into a reaction chamber at a pressure between 0.2-10 T with approximately one T being preferred, and at a temperature between 450-625°C with 550°C being preferred. An inert gas or gases such as argon, helium, and nitrogen can be included in the gas mix in order to prevent undesired tungsten-silicide deposition on the reaction chamber and to provide a carrier for WF<sub>6</sub>.

It is to be noted that DCS based tungsten-silicide layer 306 can be formed uniformly and with a smooth surface, as shown in Figure 3b, because it is formed onto a doped amorphous silicon layer as opposed to a doped polysilicon layer. Because amorphous silicon is devoid of a long range periodic structures, it has few grain boundaries. Because amorphous silicon lacks substantial grain boundaries, dopants are unable to readily diffuse to the surface where they can inhibit nucleation and therefore good film growth. Thus, by forming DCS based tungsten-silicide layer 306 directly onto amorphous silicon layer 304, a smooth and uniform tungsten-silicide layer can be grown without the need for a "capping" layer as required with doped polysilicon.

In order to prevent a tungsten rich interface, and initial nucleation step may be utilized prior to deposition of DCS based tungsten-silicide layer 308, if desired. Such a nucleation step would occur for approximately 20 seconds and would utilize the same process gases and parameters as DCS based tungsten-silicide film 306, except that the nucleation step would occur at a slightly higher pressure, approximately 1.5 T in the preferred embodiment, than the bulk deposition, which in the preferred embodiment is one T.

Additionally, after deposition of DCS based tungsten-silicide layer 306, a "monosilane cap" can be formed on DCS based tungsten-silicide layer 306, if desired. Flowing monosilane over tungsten-silicide layer 306 allows any loose tungsten bonds to react with monosilane and form tungsten-silicide. A monosilane cap can be formed as described above.

After formation of a "monosilane cap," substrate

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200 is removed from the tungsten-silicide deposition chamber. Substrate 200 is then eventually annealed to activate and drive the dopants in amorphous silicon layer 304 and to alter the stoichiochemistry of DCS based tungsten-silicide film 306 to decrease its resistance. Any well known anneal method and apparatus can be utilized to anneal substrate 200, such as a rapid thermal anneal (RTA), at approximately 1000°C, in a nitrogen ambient, for a time between 30-60 seconds, or a standard furnace anneal. It is to be appreciated that the anneal step converts amorphous silicon 304 into polycrystalline silicon.

At this time, the formation of a novel amorphous silicon/DCS based tungsten-silicide composite film 308 is now complete. The novel silicon/tungsten-silicide composite film 308 can be formed thin (less than 1000Å) and uniformly across the surface of a wafer. Very little, if any, fluorine is incorporated into a composite film 308 because tungsten-silicide layer 306 is deposited from DCS. Conventional patterning, doping, and interconnection techniques can be used to complete the fabrication of an integrated circuit.

It is to be noted that although the present invention has been described with respect to the formation of a gate electrode, the processes of the present invention can be used anywhere low resistance silicon films are needed, such as interconnects. Additionally, although the present invention has been described with respect to the formation of thin (less than 1000Å) films, the present invention can be used to form films with thicknesses greater than 1000Å. As such, the detailed description of the present invention is to be taken as illustrative rather than limiting whereby the scope of the present invention is to be measured by the appended claims which follow.

Thus, thin, low resistance silicon/tungsten-silicide composite films and their methods of fabrication have been described.

#### Claims

 A method of forming a silicide on a silicon layer comprising the steps of:

forming a first tungsten-silicide layer on said silicon layer wherein said first tungsten-silicide layer is formed from a first gas mix comprising monosilane (SiH<sub>4</sub>); and

forming a second tungsten-silicide layer on said first tungsten-silicide layer wherein said second tungsten-silicide layer is formed from a second gas mix comprising dichlorosilane (DCS).

A method as claimed in claim 1 further comprising the step of:

exposing said silicon layer to a third gas comprising monosilane (SiH<sub>4</sub>) prior to forming said first tungsten-silicide layer.

- A method as claimed in claim 1 or claim 2 further comprising the step of exposing said second tungsten-silicide layer to a fourth gas comprising silane.
  - 4. A method as claimed in any one of the preceding claims, wherein said first gas mix further comprises WF<sub>6</sub> and said first gas mix has a WF<sub>6</sub>:SiH<sub>4</sub> ratio of approximately 1:100.
- A method as claimed in any one of the preceding claims wherein said first tungsten-silicide layer is formed at a pressure between 0.2-10 T and at a temperature between 450-625°C.
- A method as claimed in any one of the preceding claims wherein said silicon layer is polycrystalline silicon.
- A method as claimed in any one of the preceding claims wherein said silicon layer is amorphous silicon.
- 25 8. A method of forming a silicide on a doped polysilicon layer comprising the steps of:

exposing said doped polysilicon layer to a first gas mix comprising monosilane (SiH<sub>4</sub>) to form a monosilane exposed polysilicon surface; forming a first tungsten-silicide layer on said monosilane exposed doped polysilicon surface wherein said first tungsten-silicide layer in

wherein said first tungsten-silicide layer is formed from a reduction reaction of SiH<sub>4</sub> and WF<sub>6</sub>; and

forming a second tungsten-silicide layer on said first tungsten-silicide layer wherein said second tungsten-silicide layer is formed from a reduction reaction of DCS and WF<sub>6</sub>.

- A method as claimed in claim 8 wherein said first tungsten-silicide layer is formed to a thickness of approximately 100Å.
- 45 10. A method as claimed in claim 8 or claim 9 wherein said first tungsten-silicide layer is formed from a gas mix having a WF<sub>6</sub>:SiH<sub>4</sub> ratio of approximately 1: 100.
- 50 11. A method as claimed in any one of the preceding claims wherein said first tungsten-silicide layer has a Si:W ratio greater than or equal to the Si:W ratio in said second tungsten-silicide layer.
- 5 12. A method as claimed in any one of claims 8 to 11 further comprising the step of exposing said second tungsten-silicide layer to a second gas mix comprising silane.

13. A composite thin film comprising:

a silicon layer; A SiH $_4$  based tungsten-silicide layer formed on said silicon layer; and a SiH $_2$ Cl $_2$  based tungsten-silicide layer formed on said SiH $_4$  based tungsten-silicide layer.

 A composite thin film as claimed in claim 13 wherein said SiH<sub>4</sub> based tungsten-silicide layer has a thickness of approximately 100Å.

15. A composite thin film as claimed in claim 13 or claim 14 wherein the Si: W ratio of said SiH<sub>4</sub> based tungsten-silicide layer is greater than or equal to the Si: W ratio of said SiH<sub>2</sub>Cl<sub>2</sub> based tungsten-silicide layer.

16. A composite thin film as claimed in any one of claims 13 to 15 wherein said silicon layer is an amorphous silicon layer.

17. A composite thin film as claimed in any one of claims 13 to 16 wherein said silicon layer is a polycrystalline silicon layer.

18. A composite thin film as claimed in any one of claims 13 to 17 wherein said polycrystalline silicon layer is doped.

19. A composite thin film as claimed in any one of claims 13 to 18 wherein said monosilane based tungsten-silicide layer is substantially thicker than said SiH<sub>2</sub>Cl<sub>2</sub> based tungsten-silicide layer.

20. A method of forming a thin film comprising the steps

forming an amorphous silicon layer; and forming a dichlorosilane based tungsten-silicide layer on said amorphous silicon layer.

 A method as claimed in claim 20 further comprising the step of annealing said thin film to convert said amorphous silicon to polysilicon.

22. A method as claimed in claim 20 or claim 21 wherein said amorphous silicon layer is doped.

23. A method as claimed in any one of claims 20 to 22 wherein said dichlorosilane based tungsten-silicide layer is formed from a reaction of WF<sub>6</sub> and SiH<sub>2</sub>Cl<sub>2</sub>.

24. A thin film comprising:

an amorphous silicon layer; and a dichlorosilane based tungsten-silicide layer on said amorphous silicon layer.

A thin film as claimed in claim 24 wherein said amorphous silicon layer is doped.

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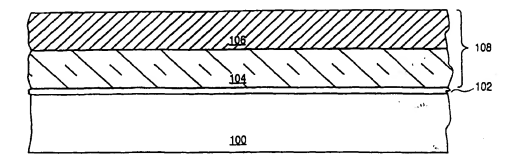


FIG. 1A

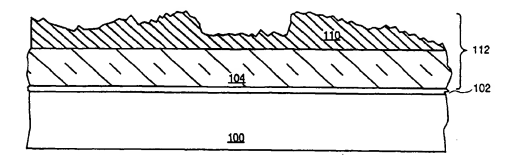


FIG. 1B

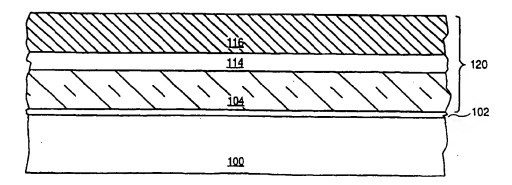
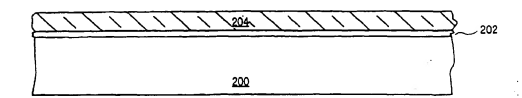
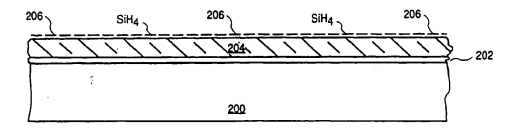


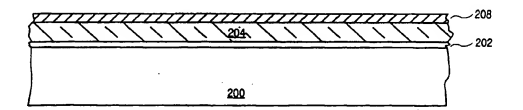
FIG. 1C



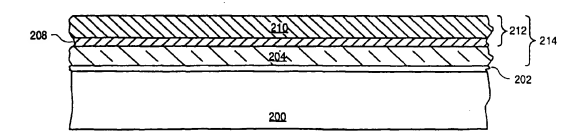
## FIG. 2A



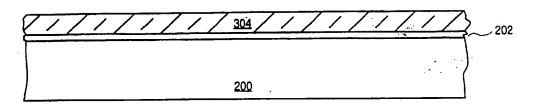
### FIG. 2B



## FIG. 2C



## FIG. 2D



# FIG. 3A

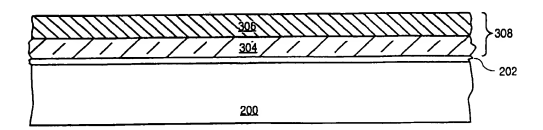


FIG. 3B